

# House File 305 - Introduced

HOUSE FILE 305  
BY COMMITTEE ON HUMAN  
RESOURCES

(SUCCESSOR TO HSB 38)

## A BILL FOR

1 An Act relating to the prescribing of biological products and  
2 making penalties applicable.  
3 BE IT ENACTED BY THE GENERAL ASSEMBLY OF THE STATE OF IOWA:

1 Section 1. Section 155A.3, Code 2017, is amended by adding  
2 the following new subsections:

3 NEW SUBSECTION. 2A. "*Biological product*" means the same as  
4 defined in 42 U.S.C. §262.

5 NEW SUBSECTION. 19A. "*Interchangeable biological product*"  
6 means either of the following:

7 a. A biological product that the United States food and  
8 drug administration has licensed and has determined meets  
9 the standards for interchangeability pursuant to 42 U.S.C.  
10 §262(k)(4).

11 b. A biological product that the United States food and  
12 drug administration has determined to be therapeutically  
13 equivalent to another biological product as set forth in the  
14 latest edition or supplement of the United States food and  
15 drug administration approved drug products with therapeutic  
16 equivalence evaluations publication.

17 Sec. 2. Section 155A.28, Code 2017, is amended to read as  
18 follows:

19 **155A.28 Label of prescription drugs — interchangeable**  
20 **biological product list.**

21 1. The label of any drug, biological product, or device sold  
22 and dispensed on the prescription of a practitioner shall be in  
23 compliance with rules adopted by the board.

24 2. The board shall maintain a link on its internet site to  
25 the current list of all biological products that the United  
26 States food and drug administration has determined to be  
27 interchangeable biological products.

28 Sec. 3. Section 155A.32, Code 2017, is amended to read as  
29 follows:

30 **155A.32 Drug product selection — restrictions.**

31 1. a. If an authorized prescriber prescribes, in  
32 writing, electronically, by facsimile, or orally, a drug  
33 by its brand or trade name, the pharmacist may exercise  
34 professional judgment in the economic interest of the patient  
35 by selecting a drug product with the same generic name

1 and demonstrated bioavailability as the one drug product  
2 prescribed for dispensing and sale to the patient. If the  
3 cost of the prescription or any part of it will be paid by  
4 expenditure of public funds authorized under [chapter 249A](#), the  
5 pharmacist shall exercise professional judgment by selecting  
6 a drug product with the same generic name and demonstrated  
7 bioavailability as the one drug product prescribed for  
8 dispensing and sale. ~~If the pharmacist exercises drug product~~  
9 ~~selection, the pharmacist shall inform the patient of the~~  
10 ~~savings which the patient will obtain as a result of the drug~~  
11 ~~product selection and pass on to the patient no less than fifty~~  
12 ~~percent of the difference in actual acquisition costs between~~  
13 ~~the drug prescribed and the drug substituted.~~

14 b. If an authorized prescriber prescribes a biological  
15 product, the pharmacist may exercise professional judgment in  
16 the economic interest of the patient by selecting a biological  
17 product that is an interchangeable biological product for the  
18 biological product prescribed for dispensing and sale to the  
19 patient. If the cost of the prescription or any part of it will  
20 be paid by expenditure of public funds authorized under chapter  
21 249A, the pharmacist shall exercise professional judgment by  
22 selecting a biological product that is an interchangeable  
23 biological product for the biological product prescribed for  
24 dispensing and sale.

25 2. The pharmacist shall not exercise the drug or biological  
26 product selection described in [this section](#) if ~~either~~ any of  
27 the following is true:

28 a. The prescriber specifically indicates that no drug or  
29 biological product selection shall be made.

30 b. The person presenting the prescription indicates that  
31 only the specific drug product prescribed should be dispensed.  
32 However, this paragraph does not apply if the cost of the  
33 prescription or any part of it will be paid by expenditure of  
34 public funds authorized under [chapter 249A](#).

35 3. If selection of a generically equivalent drug product

1 or an interchangeable biological product is made under this  
 2 section, the pharmacist making the selection shall inform the  
 3 patient and note that fact and the name of the manufacturer of  
 4 the selected drug on the prescription presented by the patient  
 5 or the patient's adult representative or transmitted by the  
 6 prescriber or the prescriber's authorized agent.

7 4. a. Within five business days following the dispensing  
 8 of a biological product, the dispensing pharmacist or the  
 9 pharmacist's designee shall make an entry of the specific  
 10 biological product provided to the patient, including the name  
 11 of the biological product and the manufacturer. The entry  
 12 shall be electronically accessible to the prescriber through  
 13 one of the following means:

14 (1) An interoperable electronic medical records system.

15 (2) An electronic prescribing technology.

16 (3) A pharmacy benefit management system.

17 (4) A pharmacy record.

18 b. An entry into an electronic records system as described  
 19 in this subsection is presumed to provide notice to the  
 20 prescriber. If the entry is not made electronically, the  
 21 pharmacist shall communicate the name and manufacturer of the  
 22 biological product dispensed to the prescriber using facsimile,  
 23 telephone, electronic transmission, or other prevailing means.

24 c. Communication under this subsection shall not be required  
 25 in either of the following circumstances:

26 (1) There is no federal food and drug  
 27 administration-approved interchangeable biological product for  
 28 the product prescribed.

29 (2) A refill prescription is not changed from the product  
 30 dispensed on the prior filling of the prescription.

#### 31 EXPLANATION

32 The inclusion of this explanation does not constitute agreement with  
 33 the explanation's substance by the members of the general assembly.

34 This bill adopts by reference to federal law a definition of  
 35 "biological product" and defines "interchangeable biological

1 product". As described by the United States food and drug  
2 administration, a "biological product" is a medical product,  
3 often made from a variety of natural sources, used for a broad  
4 range of diseases or conditions, particularly chronic, serious,  
5 or life-threatening conditions such as cancer and rheumatoid  
6 arthritis.

7 The bill provides that pharmacists may use professional  
8 judgment to distribute an interchangeable biological product  
9 when an authorized prescriber prescribes a biological product.  
10 The bill requires the board of pharmacy to maintain a link  
11 on its internet site to the current list of all biological  
12 products that the United States food and drug administration  
13 has determined to be interchangeable biological products.

14 The bill provides that a pharmacist may not dispense  
15 an interchangeable biological product if the prescriber  
16 specifically indicates that no product selection shall be made  
17 or the person presenting the prescription indicates that only  
18 the specific biological product prescribed should be dispensed.

19 The bill removes a provision that requires a pharmacist  
20 to pass on to the patient no less than fifty percent of the  
21 difference in actual acquisition costs between the drug  
22 prescribed and the drug substituted and inform a patient of  
23 those savings if the pharmacist makes a drug product selection.

24 The bill requires a pharmacist to notify a patient whenever  
25 the pharmacist selects a generically equivalent drug product or  
26 an interchangeable biological product.

27 The bill requires that within five days of dispensing an  
28 interchangeable biological product, a pharmacist must make  
29 an entry into one of a specified type of electronic records  
30 systems noting the name and manufacturer of the biological  
31 product. According to the bill, such an entry is deemed to  
32 provide notice to the prescriber if done electronically. If  
33 it is not done electronically, the pharmacist must otherwise  
34 provide the name and manufacturer of the biological product to  
35 the prescriber. Such communication is not required if a refill

1 prescription is not changed from the product dispensed on the  
2 prior filling of the prescription.

3     A person who violates these provisions with regard to  
4 a noncontrolled substance shall be guilty of a serious  
5 misdemeanor for a first violation, an aggravated misdemeanor  
6 for a second offense or if the person has been convicted with  
7 a violation of laws relating to prescription drugs or devices  
8 in other jurisdictions, or a class "D" felony for a third  
9 offense or a second offense with prior conviction in another  
10 jurisdiction. A person who violates these provisions with  
11 regard to a controlled substance shall be punished pursuant to  
12 Code section 124.401, subsection 1, and other provisions of  
13 Code chapter 124, division IV.